This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problems Mailbox.

THIS PAGE BLANK (USPTO)

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 04.03.1998 Bulletin 1998/10

(51) Int Cl.6: G03F 7/038

(21) Application number: 97306199.7

(22) Date of filing: 15.08.1997

(84) Designated Contracting States:
AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC
NL PT SE

(30) Priority: 29.08.1996 US 705365

(71) Applicant: XEROX CORPORATION
Rochester New York 14644 (US)

(72) Inventors:

Narang, Ram S.
 Fairport NY 14450 (US)

• Fuller, Timothy J. Pittsford NY 14534-4023 (US)

(74) Representative: Pike, Christopher Gerard et al Rank Xerox Ltd., Patent Department, Parkway

Marlow, Buckinghamshire SL7 1YL (GB)

(54) Hydroxyalkated high performance curable polymers

(57) Disclosed is a composition which comprises (a) a defined polymer containing at least some monomer repeat units with photosensitivity-imparting substituents which enable crosslinking or chain extension of the polymer upon exposure to actinic radiation, wherein said photosensitivity-imparting substituents are hydroxyalkyl

groups; (b) at least one member selected from the group consisting of photoinitiators and sensitizers; and (c) an optional solvent. Also disclosed are processes for preparing the above polymers and methods of preparing thermal ink jet printheads containing the above polymers.

Description

5

10

The present invention is directed to curable polymers and to photoresists and thermal ink jet printheads containing these polymers. More specifically, the present invention is directed to high performance polymers substituted with hydroxyalkyl groups.

In microelectronics applications, there is a great need for low dielectric constant, high glass transition temperature, thermally stable, photopatternable polymers for use as interlayer dielectric layers and as passivation layers which protect microelectronic circuitry. Poly(imides) are widely used to satisfy these needs; these materials, however, have disadvantageous characteristics such as relatively high water sorption and hydrolytic instability. There is thus a need for high performance polymers which can be effectively photopatterned and developed at high resolution.

One particular application for such materials is the fabrication of ink jet printheads.

Other microelectronics applications include printed circuit boards, lithographic printing processes, and interlayer dielectrics.

Copending application U.S. Serial No. 08/705,375 discloses an improved composition comprising a photopatternable polymer containing at least some monomer repeat units with photosensitivity-imparting substituents.

Copending application U.S. Serial No. 08/705,488 discloses a composition comprising a polymer with a weight average molecular weight of from about 1,000 to about 65,000, said polymer containing at least some monomer repeat units with a first, photosensitivity-imparting substituent which enables crosslinking or chain extension of the polymer upon exposure to actinic radiation, said polymer also containing a second, thermal sensitivity-imparting substituent which enables further polymerization of the polymer upon exposure to temperatures of about 140°C and higher.

Copending application U.S. Serial No. 08/697,761 discloses a process which comprises reacting a defined polymer with (i) a formaldehyde source, and (ii) an unsaturated acid in the presence of an acid catalyst, thereby forming a curable polymer with unsaturated ester groups.

Copending application U.S. Serial No. 08/705,463 discloses a process which comprises reacting a defined polymer with an acetyl halide and dimethoxymethane in the presence of a halogen-containing Lewis acid catalyst and methanol, thereby forming a haloalkylated polymer.

Copending application U.S. Serial No. 08/705,479 discloses a process which comprises reacting a haloalkylated aromatic polymer with a material selected from the group consisting of unsaturated ester salts, alkoxide salts, alkyl-carboxylate salts, and mixtures thereof, thereby forming a curable polymer having functional groups corresponding to the selected salt.

Copending application U.S. Serial No. 08/705,376 discloses a composition which comprises a mixture of (A) a first component comprising a defined polymer, at least some of the monomer repeat units of which have at least one photosensitivity-imparting group thereon, and (B) a second component which comprises either (1) a polymer having a second degree of photosensitivity-imparting group substitution or (2) a reactive diluent having at least one photosensitivity-imparting group per molecule and having a fourth degree of photosensitivity-imparting group substitution.

Copending application U.S. Serial No. 08/705,372 discloses a composition which comprises a defined polymer containing at least some monomer repeat units with photosensitivity-imparting substituents which enable crosslinking or chain extension of the polymer upon exposure to actinic radiation wherein said photosensitivity-imparting substituents are allyl ether groups, epoxy groups, or mixtures thereof.

Copending application U.S. Serial No. 08/697,760 discloses a composition which comprises a defined polymer containing at least some monomer repeat units with water-solubility-imparting substituents and at least some monomer repeat units with photosensitivity-imparting substituents which enable crosslinking or chain extension of the polymer upon exposure to actinic radiation.

While known compositions and processes are suitable for their intended purposes, a need remains for improved materials suitable for microelectronics applications. A need also remains for improved ink jet printheads. Further, there is a need for photopatternable polymeric materials which are heat stable, electrically insulating, and mechanically robust. Additionally, there is a need for photopatternable polymeric materials which are chemically inert with respect to the materials that might be employed in ink jet ink compositions. There is also a need for photopatternable polymeric materials which exhibit low shrinkage during post-cure steps in microelectronic device fabrication processes. In addition, a need remains for photopatternable polymeric materials which exhibit a relatively long shelf life. Further, there is a need for photopatternable polymeric materials which, in the cured form, exhibit good solvent resistance. There is also a need for photopatternable polymeric materials which, when applied to microelectronic devices by spin casting techniques and cured, exhibit reduced edge bead and no apparent lips and dips. In addition, there remains a need for processes for preparing photopatternable polymeric materials with the above advantages. Further, a need remains for processes for preparing photopatternable polymeric materials with high aspect ratios at high resolutions by the incorporation of polymerizable groups and/or cross-linking sites pendant to the polymers. Additionally, there is a need for processes for preparing aromatic polymers having unsaturated ester functional groups

pendant to the polymer chains. There is also a need for processes for preparing photopatternable polymers having unsaturated ester functional groups pendant to the polymer chains. In addition, a need remains for photoresist materials which can be patterned as thick 30 micron films which are resistant to alkaline media. Further, there is a need for photoresist materials which can serve as interlayer dielectrics at high temperatures. Additionally, a need remains for photoresist materials which offer the advantage of no HCI liberation during thermal cure. In addition, there remains a need for photopatternable polymeric materials which have relatively low dielectric constants. Further, there is a need for photopatternable polymeric materials which exhibit reduced water sorption. Additionally, a need remains for photopatternable polymeric materials which exhibit improved hydrolytic stability, especially upon exposure to alkaline solutions. A need also remains for photopatternable polymeric materials which are stable at high temperatures, typically greater than about 150°C. There is also a need for photopatternable polymeric materials which either have high glass transition temperatures or are sufficiently crosslinked that there are no low temperature phase transitions subsequent to photoexposure. Further, a need remains for photopatternable polymeric materials with low coefficients of thermal expansion. There is a need for polymers which are thermally stable, patternable as thick films of about 30 microns or more, exhibit low T_q prior to photoexposure, have low dielectric constants, are low in water absorption, have low coefficients of expansion, have desirable mechanical and adhesive characteristics, and are generally desirable for interlayer dielectric applications, including those at high temperatures, which are also photopatternable. There is also a need for photoresist compositions with good to excellent processing characteristics.

According to one aspect of the present invention, there is provided a composition which comprises (a) a polymer containing at least some monomer repeat units with photosensitivity-imparting substituents which enable crosslinking or chain extension of the polymer upon exposure to actinic radiation, said polymer being of the formula

or

35

10

15

20

25

30

40

45

wherein x is an integer of 0 or 1, A is

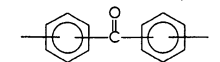
Wilesell X is an integer of 0 of 1, A is

55

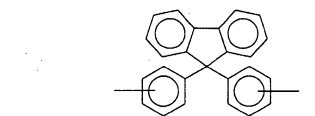
۰0.

or mixtures thereof, B is

F₃C, CF₃







wherein ${\bf v}$ is an integer of from 1 to about 20,

wherein z is an integer of from 2 to about 20,

wherein u is an integer of from 1 to about 20,

15 CH₃

30

⁵⁵ CH₃ CH₃ CH₃ CH₃ CH₃

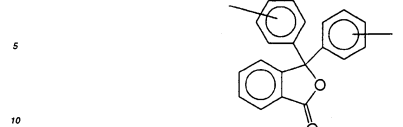
CH₃ (CH₂)w

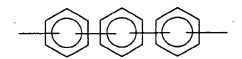
wherein w is an integer of from 1 to about 20,

40

50

55 CH₃ CH₃





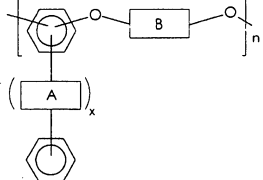


or mixtures thereof, and n is an integer representing the number of repeating monomer units, wherein said photosensitivity-imparting substituents are hydroxyalkyl groups; (b) at least one member selected from the group consisting of photoinitiators and sensitizers; and (c) an optional solvent.

According to another aspect of the present invention, there is provided a process which comprises the steps of: (a) reacting a haloalkylated polymer of the formula

В

40 or 45



30

15

20

25

35

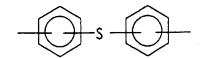
50

wherein x is an integer of 0 or 1, A is

-0-,

-C(CH₃)₂-,

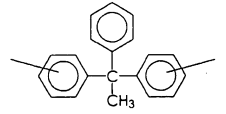
or mixtures thereof, B is



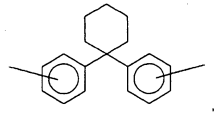
wherein v is an integer of from 1 to about 20,

wherein z is an integer of from 2 to about 20,

wherein u is an integer of from 1 to about 20,







wherein w is an integer of from 1 to about 20,

or mixtures thereof, and n is an integer representing the number of repeating monomer units, with water and a base, thereby forming a polymer with hydroxyalkyl groups; and optionally (b) converting the hydroxyalkyl groups to unsaturated ester groups.

Figure 1 is an enlarged schematic isometric view of an example of a printhead mounted on a daughter board

showing the droplet emitting nozzles.

Figure 2 is an enlarged cross-sectional view of Figure 1 as viewed along the line 2-2 thereof and showing the electrode passivation and ink flow path between the manifold and the ink channels.

Figure 3 is an enlarged cross-sectional view of an alternate embodiment of the printhead in Figure 1 as viewed along the line 2-2 thereof.

The present invention is directed to curable polymers having hydroxyalkyl functional groups. The polymers are of the following formula:

10

5

A X O B

or

20

15

20

25

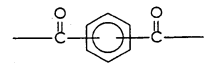
30

B O B

wherein x is an integer of 0 or 1, A is

40

35



45

50

. .

or mixtures thereof, B is

-O-,

-C(CH₃)₂-,

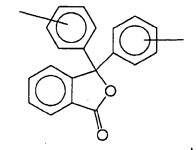
wherein v is an integer of from 1 to about 20, and preferably from 1 to about 10,

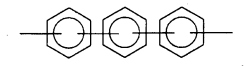
(CH₂)√

wherein z is an integer of from 2 to about 20, and preferably from 2 to about 10,

wherein u is an integer of from 1 to about 20, and preferably from 1 to about 10,

wherein w is an integer of from 1 to about 20, and preferably from 1 to about 10, $\,$





other similar bisphenol derivatives, or mixtures thereof, and n is an integer representing the number of repeating mon-

10

15

20

25

30

35

40

45

50

omer units. The value of n is such that the weight average molecular weight of the material is from about 1,000 to about 100,000, preferably from about 1,000 to about 65,000, more preferably from about 1,000 to about 40,000, and even more preferably from about 3,000 to about 25,000, although the weight average molecular weight can be outside these ranges. Preferably, n is an integer of from about 2 to about 70, more preferably from about 5 to about 70, and even more preferably from about 8 to about 50. The phenyl groups and the A and/or B groups may also be substituted, although the presence of two or more substituents on the B group ortho to the oxygen groups can render substitution difficult. Substituents can be present on the polymer either prior to or subsequent to the placement of photosensitivityimparting functional groups thereon. Substituents can also be placed on the polymer during the process of placement of photosensitivity-imparting functional groups thereon. Examples of suitable substituents include (but are not limited to) alkyl groups, including saturated, unsaturated, and cyclic alkyl groups, preferably with from 1 to about 6 carbon atoms, substituted alkyl groups, including saturated, unsaturated, and cyclic substituted alkyl groups, preferably with from 1 to about 6 carbon atoms, aryl groups, preferably with from 6 to about 24 carbon atoms, substituted aryl groups, preferably with from 6 to about 24 carbon atoms, arylalkyl groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyl groups, preferably with from 7 to about 30 carbon atoms, alkoxy groups, preferably with from 1 to about 6 carbon atoms, substituted alkoxy groups, preferably with from 1 to about 6 carbon atoms, aryloxy groups, preferably with from 6 to about 24 carbon atoms, substituted aryloxy groups, preferably with from 6 to about 24 carbon atoms, arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, hydroxy groups, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, ester groups, amide groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, mercapto groups, nitroso groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, and the like, wherein the substituents on the substituted alkyl groups, substituted aryl groups, substituted arylalkyl groups, substituted alkoxy groups, substituted aryloxy groups, and substituted arylalkyloxy groups can be (but are not limited to) hydroxy groups, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, aldehyde groups, ketone groups, ester groups, amide groups, carboxylic acid groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, cyano groups, nitrile groups, mercapto groups, nitroso groups, halogen atoms, nitro groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, mixtures thereof, and the like, wherein two or more substituents can be joined together to form a ring. Processes for the preparation of these materials are known, and disclosed in, for example, P. M. Hergenrother, J. Macromol. Sci. Rev. Macromol. Chem., C19 (1), 1-34 (1980); P. M. Hergenrother,

B. J. Jensen, and S. J. Havens, Polymer, 29, 358 (1988); B. J. Jensen and P.M. Hergenrother, "High Performance Polymers," Vol. 1, No. 1) page 31 (1989), "Effect of Molecular Weight on Poly(arylene ether ketone) Properties"; V. Percec and B. C. Auman, Makromol. Chem. 185, 2319 (1984); "High Molecular Weight Polymers by Nickel Coupling of Aryl Polychlorides," I. Colon, G. T. Kwaiatkowski, J. of Polymer Science, Part A, Polymer Chemistry, 28, 367 (1990); M. Ueda and T. Ito, Polymer J., 23 (4), 297 (1991); "Ethynyl-Terminated Polyarylates: Synthesis and Characterization, S. J. Havens and P. M. Hergenrother, J. of Polymer Science: Polymer Chemistry Edition, 22, 3011 (1984); "Ethynyl-Terminated Polysulfones: Synthesis and Characterization, P. M. Hergenrother, J. of Polymer Science: Polymer Chemistry Edition, 20, 3131 (1982); K. E. Dukes, M. D. Forbes, A. S. Jeevarajan, A. M. Belu, J. M. DeDimone, R. W. Linton, and V. V. Sheares, Macromolecules, 29, 3081 (1996); G. Hougham, G. Tesoro, and J. Shaw, Polym. Mater. Sci. Eng., 61, 369 (1989); V. Percec and B. C. Auman, Makromol. Chem, 185, 617 (1984); "Synthesis and characterization of New Fluorescent Poly(arylene ethers), "S. Matsuo, N. Yakoh, S. Chino, M. Mitani, and S. Tagami, Journal of Polymer Science: Part A: Polymer Chemistry, 32, 1071 (1994); "Synthesis of a Novel Naphthalene-Based Poly(arylene ether ketone) with High Solubility and Thermal Stability," Mami Ohno, Toshikazu Takata, and Takeshi Endo, Macromolecules, 27, 3447 (1994); "Synthesis and Characterization of New Aromatic Poly(ether ketones)," F. W. Mercer, M. T. Mckenzie, G. Merlino, and M. M. Fone, J. of Applied Polymer Science, 56, 1397 (1995); H. C. Zhang, T. L. Chen, Y. G. Yuan, Chinese Patent CN 85108751 (1991); "Static and laser light scattering study of novel thermoplastics. 1. Phenolphthalein poly(aryl ether ketone), "C. Wu, S. Bo, M. Siddiq, G. Yang and T. Chen, Macromolecules, 29, 2989 (1996); "Synthesis of t-Butyl-Substituted Poly(ether ketone) by Nickel-Catalyzed Coupling Polymerization of Aromatic Dichloride", M. Ueda, Y. Seino, Y. Haneda, M. Yoneda, and J.-I. Sugiyama, Journal of Polymer Science: Part A: Polymer Chemistry, 32, 675 (1994); "Reaction Mechanisms: Comb-Like Polymers and Graft Copolymers from Macromers 2. Synthesis, Characterzation and Homopolymerization of a Styrene Macromer of Poly(2,6-dimethyl-1,4-phenylene Oxide), "V. Percec, P. L. Rinaldi, and B. C. Auman, Polymer Bulletin, 10, 397 (1983); Handbook of Polymer Synthesis Part A, Hans R. Kricheldorf, ed., Marcel Dekker, Inc., New York-Basel-Hong Kong (1992); and "Introduction of Carboxyl Groups into Crosslinked Polystyrene, "C. R. Harrison, P. Hodge, J. Kemp, and G. M. Perry, Die Makromolekulare Chemie, 176, 267 (1975).

For applications wherein the photopatternable polymer is to be used as a layer in a thermal ink jet printhead, the polymer preferably has a number average molecular weight of from about 3,000 to about 20,000, more preferably from about 3,000 to about 10,000, and even more preferably from about 3,500 to about 6,500.

The polymer can be hydroxyalkylated at one or more sites, as follows:

A B O B O P

A X B O B O n

50 or

5

25

30

35

40

45

55

or

5

10

15 В ŔOH

20

25

30

45

50

55

wherein R is an alkyl group, including both saturated, unsaturated, linear, branched, and cyclic alkyl groups, preferably with from 1 to about 11 carbon atoms, more preferably with from 1 to about 5 carbon atoms, even more preferably with from 1 to about 3 carbon atoms, and most preferably with 1 carbon atom, or a substituted alkyl group, preferably with from 1 to about 11 carbon atoms, more preferably with from 1 to about 5 carbon atoms, even more preferably with from 1 to about 3 carbon atoms, and most preferably with 1 carbon atom, and X is a halogen atom, such as fluorine, chlorine, bromine, or iodine. Examples of suitable substituents on the substituted alkyl group include (but are not limited to) alkyl groups, including saturated, unsaturated, linear, branched, and cyclic alkyl groups, preferably with from 1 to about 6 carbon atoms, substituted alkyl groups, preferably with from 1 to about 6 carbon atoms, aryl groups, preferably with from 6 to about 24 carbon atoms, substituted aryl groups, preferably with from 6 to about 24 carbon atoms, arylalkyl groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyl groups, preferably with from 7 to about 30 carbon atoms, alkoxy groups, preferably with from 1 to about 6 carbon atoms, substituted alkoxy groups, preferably with from 1 to about 6 carbon atoms, aryloxy groups, preferably with from 6 to about 24 carbon atoms, substituted aryloxy groups, preferably with from 6 to about 24 carbon atoms, arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, ester groups, amide groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, mercapto groups, nitroso groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, and the like, wherein the substituents on the substituted alkyl groups, substituted aryl groups, substituted arylalkyl groups, substituted alkoxy groups, substituted aryloxy groups, and substituted arylalkyloxy groups can be (but are not limited to) hydroxy groups, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, aldehyde groups, ketone groups, ester groups, amide groups, carboxylic acid groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, cyano groups, nitrile groups, mercapto groups, nitroso groups, halogen atoms, nitro groups, sulfone groups, acyl groups, acid anhydride groups, and azide groups, mixtures thereof, wherein two or more substituents can be joined together to form a ring.

The hydroxymethylation of a polymer of the above formula can be accomplished by reacting the polymer in solution with formaldehyde or paraformaldehyde and a base, such as sodium hydroxide, potassium hydroxide, calcium hydroxide, ammonium hydroxide, tetramethylammonium hydroxide, or the like. The polymer is dissolved in a suitable solvent, such as 1,1,2,2-tetrachloroethane or the like, and is allowed to react with the formaldehyde or paraformaldehyde. Examples of solvents suitable for the reaction include 1,1,2,2-tetrachloroethane, as well as methylene chloride, provided a suitable pressure reactor is used. Typically, the reactants are present in relative amounts by weight of about 44.5 parts paraformaldehyde or 37 parts formaldehyde, about 1 part base, about 200 parts 1,1,2,2-tetrachloroethane, and about 100 parts polymer.

The general reaction scheme is as follows:

wherein a, b, c, and d are each integers of 0, 1, 2, 3, or 4, provided that at least one of a, b, c, and d is equal to or greater than 1 in at least some of the monomer repeat units of the polymer, and n is an integer representing the number of repeating monomer units. Substitution is generally random, although the substituent often indicates a preference for the B group, and a particular preference for the sites ortho to oxygen on the B group, and any given monomer repeat unit may have no hydroxymethyl substituents, one hydroxymethyl substituent, or two or more hydroxymethyl substituents. Most commonly, each aromatic ring will have either no hydroxymethyl groups or one hydroxymethyl group.

Typical reaction temperatures are from about 50 to about 125°C, and preferably from about 85 to about 110°C, although the temperature can be outside these ranges. Typical reaction times are from about 4 to about 24 hours, and preferably from about 4 to about 6 hours, although the time can be outside these ranges. Longer reaction times generally result in higher degrees of hydroxymethylation. Different degrees of hydroxymethylation may be desirable for different applications. Too high a degree of substitution may lead to excessive sensitivity, resulting in crosslinking or chain extension of both exposed and unexposed polymer material when the material is exposed imagewise to activating radiation, whereas too low a degree of substitution may be undesirable because of resulting unnecessarily long exposure times or unnecessarily high exposure energies. For applications wherein the photopatternable polymer is to be used as a layer in a thermal ink jet printhead, the degree of substitution (i.e., the average number of hydroxymethyl groups per monomer repeat unit) preferably is from about 0.25 to about 2.0, and more preferably from about 0.5 to about 0.8, although the degree of substitution can be outside these ranges for ink jet printhead applications. This amount of substitution corresponds to from about 0.8 to about 1.3 milliequivalents of hydroxymethyl per gram of resin.

Polymers of the above formula can also be hydroxyalkylated by first preparing the haloalkylated derivative and then replacing at least some of the haloalkyl groups with hydroxyalkyl groups. For example, the haloalkylated polymer can be hydroxyalkylated by alkaline hydrolysis of the haloalkylated polymer. The hydroxy groups replace the halide atoms in the haloalkyl groups on the polymer; accordingly, the number of carbon atoms in the haloalkyl group generally corresponds to the number of carbon atoms in the hydroxyalkyl group. Examples of suitable reactants include sodium hydroxide, potassium hydroxide, calcium hydroxide, ammonium hydroxide, tetraalkyl ammonium hydroxides, such as tetrabutyl ammonium hydroxide. Examples of solvents suitable for the reaction include 1,1,2,2-tetrachloroethane, methylene chloride, and water. Typically, the reactants are present in relative amounts with respect to each other by weight of about 13.8 parts haloalkylated polymer, about 50 parts solvent, and about 30.6 parts base (containing 23 parts tetrabutylammonium hydroxide in water). After a clear solution is obtained, 30 milliliters of sodium hydroxide (50 percent aqueous solution) is added. After 16 hours at about 25°C, the organic layer is washed with water, dried over magnesium sulfate, and poured into methanol (1 gallon) to precipitate the polymer.

The general reaction scheme, illustrated below for the chloromethylated polymer, is as follows:

5
$$(CH_{2}CI)_{0} \qquad (CH_{2}CI)_{0} \qquad (CH_{2}CI)_{0}$$

$$H_{2}O \qquad OH^{\Theta}$$

$$(CH_{2}OH)_{f} \qquad (CH_{2}OH)_{g} \qquad (CH_{2}OH)_{h}$$

$$(CH_{2}OH)_{e} \qquad (CH_{2}OH)_{g} \qquad (CH_{2}OH)_{h}$$

$$(CH_{2}OH)_{e} \qquad (CH_{2}OH)_{g} \qquad (CH_{2}OH)_{h}$$

$$(CH_{2}OH)_{e} \qquad (CH_{2}OH)_{g} \qquad (CH_{2}OH)_{h}$$

35

40

45

50

wherein a, b, c, d, e, f, g, h, i, j, k, and m are each integers of 0, 1, 2, 3, or 4, provided that the sum of i+e is no greater than 4, the sum of j+f is no greater than 4, the sum of k+g is no greater than 4, and the sum of m+h is no greater than 4, provided that at least one of a, b, c, and d is equal to or greater than 1 in at least some of the monomer repeat units of the polymer, and provided that at least one of e, f, g, and h is equal to at least 1 in at least some of the monomer repeat units of the polymer, and n is an integer representing the number of repeating monomer units.

Higher degrees of haloalkylation generally enable higher degrees of substitution with hydroxyalkyl groups, and thereby enable greater photosensitivity of the polymer. Different degrees of substitution may be desirable for different applications. Too high a degree of substitution may lead to excessive sensitivity, resulting in crosslinking or chain extension of both exposed and unexposed polymer material when the material is exposed imagewise to activating radiation, whereas too low a degree of substitution may be undesirable because of resulting unnecessarily long exposure times or unnecessarily high exposure energies. For applications wherein the photopatternable polymer is to be used as a layer in a thermal ink jet printhead, the degree of substitution (i.e., the average number of hydroxyalkyl groups per monomer repeat unit) preferably is from about 0.25 to about 2.0, and more preferably from about 0.5 to about 0.8, although the degree of substitution can be outside these ranges for ink jet printhead applications. Optimum amounts of substitution are from about 0.8 to about 1.3 milliequivalents of hydroxyalkyl group per gram of resin.

Some or all of the haloalkyl groups can be replaced with hydroxyalkyl substituents. Longer reaction times generally lead to greater degrees of substitution of haloalkyl groups with hydroxyalkyl substituents.

Typical reaction temperatures are from about 25 to about 120°C, and preferably from about 25 to about 50°C, although the temperature can be outside this range. Typical reaction times are from about 1 to about 24 hours, and preferably from about 10 to about 16 hours, although the time can be outside these ranges.

The polymer to be substituted can be haloalkylated by any desired or suitable process. For example, suitable processes for haloalkylating polymers include reaction of the polymers with formaldehyde and hydrohalic acid, bishalomethyl ether, halomethyl methyl ether, octylhalomethyl ether, or the like, generally in the presence of a Lewis acid catalyst. Bromination of a methyl group on the polymer can also be accomplished with elemental bromine via a free radical process initiated by, for example, a peroxide initiator or light. Halogen atoms can be substituted for other halogens already on a halomethyl group by, for example, reaction with the appropriate hydrohalic acid or halide salt. Methods for the halomethylation of polymers are also disclosed in, for example, "Chloromethylation of Condensation Polymers Containing an oxy-1,4-phenylene Backbone," W. H. Daly et al., *Polymer Preprints*, Vol. 20, No. 1, 835 (1979), the disclosure of which is totally incorporated herein by reference.

Haloalkylation of the polymer can be accomplished by reacting the polymer with an acetyl halide and dimethoxymethane in the presence of a halogen-containing Lewis acid catalyst such as those of the general formula

wherein n is an integer of 1, 2, 3, 4, or 5, M represents a boron atom or a metal atom, such as tin, aluminum, zinc, antimony, iron (III), gallium, indium, arsenic, mercury, copper, platinum, palladium, or the like, and X represents a halogen atom, such as fluorine, chlorine, bromine, or iodine, with specific examples including SnCl₄, AlCl₃, ZnCl₂, AlBr₃, SbF₅, Fel₃, GaBr₃, InCl₃, Asl₅, HgBr₂, CuCl, PdCl₂, or PtBr₂. A suitable haloalkylation process is described in US Serial No. 08/765,463.

5

10

20

25

30

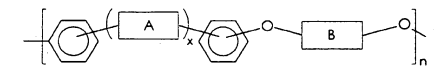
35

40

45

50

In some instances, the terminal groups on the polymer can be selected by the stoichiometry of the polymer synthesis. For example, when a polymer is prepared by the reaction of 4,4'-dichlorobenzophenone and bis-phenol A in the presence of potassium carbonate in N,N-dimethylacetamide, if the bis-phenol A is present in about 7.5 to 8 mole percent excess, the resulting polymer generally is bis-phenol A-terminated (wherein the bis-phenol A moiety may or may not have one or more hydroxy groups thereon), and the resulting polymer typically has a polydispersity (M_n/M_n) of from about 2 to about 3.5. When the bis-phenol A-terminated polymer is subjected to further reactions to place functional groups thereon, such as haloalkyl groups, and/or to convert one kind of functional group, such as a haloalkyl group, to another kind of functional group, such as an unsaturated ester group, the polydispersity of the polymer can rise to the range of from about 4 to about 6. In contrast, if the 4,4'-dichlorobenzophenone is present in about 7.5 to 8 mole percent excess, the reaction time is approximately half that required for the bis-phenol A excess reaction, the resulting polymer generally is benzophenone-terminated (wherein the benzophenone moiety may or may not have one or more chlorine atoms thereon), and the resulting polymer typically has a polydispersity of from about 2 to about 3.5. When the benzophenone-terminated polymer is subjected to further reactions to place functional groups thereon, such as haloalkyl groups, and/or to convert one kind of functional group, such as a haloalkyl group, to another kind of functional group, such as a hydroxymethyl group, the polydispersity of the polymer typically remains in the range of from about 2 to about 3.5. Similarly, when a polymer is prepared by the reaction of 4,4'-difluorobenzophenone with either 9,9'-bis (4-hydroxyphenyl)fluorene or bis-phenol A in the presence of potassium carbonate in N,N-dimethylacetamide, if the 4,4'-diffuorobenzophenone reactant is present in excess, the resulting polymer generally has benzophenone terminal groups (which may or may not have one or more fluorine atoms thereon). The well-known Carothers equation can be employed to calculate the stoichiometric offset required to obtain the desired molecular weight. (See, for example, William H. Carothers, "An Introduction to the General Theory of Condensation Polymers," Chem. Rev., 8, 353 (1931) and J. Amer. Chem. Soc., 51, 2548 (1929); see also P. J. Flory, Principles of Polymer Chemistry, Cornell University Press, Ithaca, New York (1953); the disclosures of each of which are totally incorporated herein by reference.) More generally speaking, during the preparation of polymers of the formula



the stoichiometry of the polymer synthesis reaction can be adjusted so that the end groups of the polymer are derived from the "A" groups or derived from the "B" groups. Specific functional groups can also be present on these terminal "A" groups or "B" groups, such as ethynyl groups or other thermally sensitive groups, hydroxy groups which are attached to the aromatic ring on an "A" or "B" group to form a phenolic moiety, halogen atoms which are attached to the "A" or "B" group, or the like.

Polymers with end groups derived from the "A" group, such as benzophenone groups or halogenated benzophenone groups, may be preferred for some applications because both the syntheses and some of the reactions of these materials to place substituents thereon may be easier to control and may yield better results with respect to, for example, cost, molecular weight, molecular weight range, and polydispersity (M_w/M_n) compared to polymers with end groups derived from the "B" group, such as bis-phenol A groups (having one or more hydroxy groups on the aromatic rings thereof) or other phenolic groups. While not being limited to any particular theory, it is believed that the haloalkylation reaction in particular proceeds most rapidly on the phenolic tails when the polymer is bis-phenol A terminated. Moreover, it is believed that halomethylated groups on phenolic-terminated polymers may be particularly reactive to subsequent crosslinking or chain extension. In contrast, it is generally believed that halomethylation does not take place on the terminal aromatic groups with electron withdrawing substituents, such as benzophenone, halogenated benzophenone, or the like. The "A" group terminated materials may also function as an adhesive, and in applications such as thermal ink jet printheads, the use of the crosslinked "A" group terminated polymer may reduce or eliminate the need for an epoxy adhesive to bond the heater plate to the channel plate.

The photopatternable polymer can be cured by uniform exposure to actinic radiation at wavelengths and/or energy levels capable of causing crosslinking or chain extension of the polymer through the photosensitivity-imparting groups. Alternatively, the photopatternable polymer is developed by imagewise exposure of the material to radiation at a wavelength and/or at an energy level to which the photosensitivity-imparting groups are sensitive. Typically, a photoresist composition will contain the photopatternable polymer, an optional solvent for the photopatternable polymer, an optional sensitizer, and an optional photoinitiator. Solvents may be particularly desirable when the uncrosslinked photopatternable polymer has a high T_g. The solvent and photopatternable polymer typically are present in relative amounts of from 0 to about 99 percent by weight solvent and from about 1 to 100 percent polymer, preferably are present in relative amounts of from about 40 to about 80 percent by weight polymer, and more preferably are present in relative amounts of from about 40 to about 70 percent by weight polymer.

Sensitizers absorb light energy and facilitate the transfer of energy to unsaturated bonds which can then react to crosslink or chain extend the resin. Sensitizers frequently expand the useful energy wavelength range for photoexposure, and typically are aromatic light absorbing chromophores. Sensitizers can also lead to the formation of photoinitiators, which can be free radical or ionic. When present, the optional sensitizer and the photopatternable polymer typically are present in relative amounts of from about 0.1 to about 20 percent by weight sensitizer and from about 80 to about 99.9 percent by weight photopatternable polymer, and preferably are present in relative amounts of from about 1 to about 10 percent by weight sensitizer and from about 90 to about 99 percent by weight photopatternable polymer, although the relative amounts can be outside these ranges.

Photoinitiators generally generate ions or free radicals which initiate polymerization upon exposure to actinic radiation. When present, the optional photoinitiator and the photopatternable polymer typically are present in relative amounts of from about 0.1 to about 20 percent by weight photoinitiator and from about 80 to about 99.9 percent by weight photopatternable polymer, and preferably are present in relative amounts of from about 1 to about 10 percent by weight photoinitiator and from about 90 to about 99 percent by weight photopatternable polymer, although the relative amounts can be outside these ranges.

A single material can also function as both a sensitizer and a photoinitiator.

Examples of specific sensitizers and photoinitiators include Michler's ketone (Aldrich Chemical Co.), Darocure 1173, Darocure 4265, Irgacure 184, Irgacure 261, and Irgacure 907 (available from Ciba-Geigy, Ardsley, New York), and mixtures thereof. Further background material on initiators is disclosed in, for example, Ober et al., *J.M.S. - Pure Appl. Chem.*, **A30** (12), 877-897 (1993); G. E. Green, B. P. Stark, and S. A. Zahir, "Photocrosslinkable Resin Systems, " J. *Macro. Sci. -- Revs. Macro. Chem.*, C21(2), 187 (1981); H. F. Gruber, "Photoinitiators for Free Radical Polymerization," *Prog. Polym. Sci.*, Vol. 17, 953 (1992); Johann G. Kloosterboer, "Network Formation by Chain Crosslinking Photopolymerization and Its Applications in Electronics," *Advances in Polymer Science*, <u>89</u>, Springer-Verlag Berlin Heidelberg (1988); and "Diaryliodonium Salts as Thermal Initiators of Cationic Polymerization," J. V. Crivello, T.P. Lockhart, and J. L. Lee, *J. of Polymer Science: Polymer Chemistry Edition*, <u>21</u>, 97 (1983). Sensitizers are available from, for example, Aldrich Chemical Co., Milwaukee, WI, and Pfaltz and Bauer, Waterberry, CT. Benzophenone and its derivatives can function as photosensitizers. Triphenylsulfonium and diphenyl iodonium salts are examples of typical cationic photoinitiators.

Inhibitors may also optionally be present in the photoresist containing the photopatternable polymer. Examples of suitable inhibitors include MEHQ, a methyl ether of hydroquinone, of the formula

t-butylcatechol, of the formula

5

20

25

45

50

55

hydroquinone, of the formula

and the like, the inhibitor typically present in an amount of from about 500 to about 1,500 parts per million by weight of a photoresist solution containing about 40 percent by weight polymer solids, although the amount can be outside this range.

One specific example of a class of sensitizers or initiators suitable for the polymers of the present invention is that of bis(azides), of the general formula

$$N_3$$
 R N_3 N_3

wherein R is

-R₁C=CR₂-

30 or

35

40

45

5

10

15

20

R₃ —C—

wherein R₁, R₂, R₃, and R₄ each, independently of the others, is a hydrogen atom, an alkyl group, including saturated, unsaturated, and cyclic alkyl groups, preferably with from 1 to about 30 carbon atoms, and more preferably with from 1 to about 6 carbon atoms, a substituted alkyl group, an aryl group, preferably with from 6 to about 18 carbon atoms, and more preferably with about 6 carbon atoms, a substituted aryl group, an arylalkyl group, preferably with from 7 to about 48 carbon atoms, and more preferably with from about 7 to about 8 carbon atoms, or a substituted arylalkyl group, and x is 0 or 1, wherein the substituents on the substituted alkyl, aryl, and aryl groups can be (but are not limited to) alkyl groups, including saturated, unsaturated, linear, branched, and cyclic alkyl groups, preferably with from 1 to about 6 carbon atoms, substituted alkyl groups, preferably with from 1 to about 6 carbon atoms, aryl groups, preferably with from 6 to about 24 carbon atoms, substituted anyl groups, preferably with from 6 to about 24 carbon atoms, anylalkyl groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyl groups, preferably with from 7 to about 30 carbon atoms, alkoxy groups, preferably with from 1 to about 6 carbon atoms, substituted alkoxy groups, preferably with from 1 to about 6 carbon atoms, aryloxy groups, preferably with from 6 to about 24 carbon atoms, substituted aryloxy groups, preferably with from 6 to about 24 carbon atoms, arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, substituted anylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, ester groups, amide groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, mercapto groups, nitroso groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, and the like, wherein the substituents on the substituted alkyl groups, substituted aryl groups, substituted arylalkyl groups, substituted alkoxy groups, substituted aryloxy groups, and substituted arylalkyloxy

groups can be (but are not limited to) hydroxy groups, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, aldehyde groups, ketone groups, ester groups, amide groups, carboxylic acid groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, cyano groups, nitrile groups, mercapto groups, nitroso groups, halogen atoms, nitro groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, mixtures thereof, and the like, wherein any two or more substituents can be joined together to form a ring. Examples of suitable bis(azides) include 4,4'-diazidostilbene, of the formula

$$N_3$$
—CH=CH- $\langle O \rangle$ — N_3

4,4'-diazidobenzophenone, of the formula

5

10

15

20

35

45

50

55

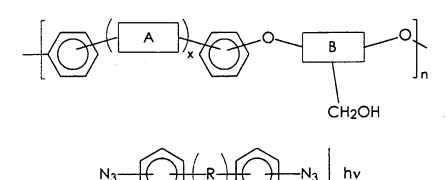
$$N_3$$
 N_3 N_3 N_3

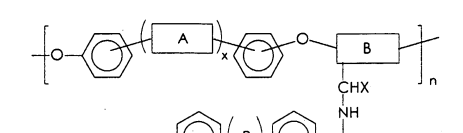
2,6-di-(4'-azidobenzal)-4-methylcyclohexanone, of the formula

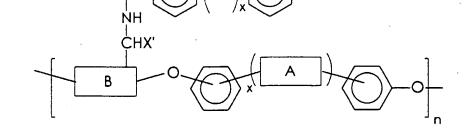
4,4'-diazidobenzalacetone, of the formula

$$N_3$$
—CH=CH-C-CH=CH—N₃

and the like. While not being limited to any particular theory, it is believed that exposure to, for example, ultraviolet radiation enables curing, as illustrated below for the hydroxymethylated polymer:







5

$$A \rightarrow H_2O$$

10

 $A \rightarrow H_2O$

15

 CH_2

20

 CH_2

A

 $A \rightarrow A$
 CH_2

27

 CH_2

28

 CH_2

A

 CH_2

wherein X and X' each, independently of the other, is -H or -OH.

55

Alternatively, a hydroxyalkylated polymer can be further reacted to render it more photosensitive. For example, a hydroxymethylated polymer of the formula

can react with an unsaturated ester isocyanate, typically of the general formula

5

10

15

20

25

30

35

40

45

50

wherein R_1 is an unsaturated alkyl group (which may be cyclic, branched, or linear), typically with from 1 to about 11 carbon atoms, or an unsaturated arylalkyl group, typically with from 7 to about 18 carbon atoms, and R_2 is an alkyl group (and may be either saturated or unsaturated, cyclic, branched, or linear), typically with from 1 to about 11 carbon atoms, or an arylalkyl group (and may be either saturated or unsaturated, cyclic, branched, or linear), typically with from 7 to about 18 carbon atoms, with specific examples including isocyanato ethyl acrylate, isocyanato ethyl cinnamate, isocyanato-ethyl methacrylate, of the formula

(available from Polysciences, Warrington, PA), other isocyanato alkyl unsaturated esters, or the like. A photoactive polymer is then formed, of, for the specific example with isocyanato ethyl methacrylate, the formula

This reaction can be carried out in methylene chloride at 25°C with 1 part by weight polymer, 1 part by weight isocyanatoethyl methacrylate, and 50 parts by weight methylene chloride. Typical reaction temperatures are from about 0 to about 50°C, with 10 to 25°C preferred. Typical reaction times are between about 1 and about 24 hours, with about 16 hours preferred. While not being limited to any particular theory, it is believed that during exposure to, for example, ultraviolet radiation, the ethylenic bond opens and crosslinking or chain extension occurs at that site. The crosslinks or chain extensions formed are believed to be via groups of the general formula

15

20

25

30

35

5

10

wherein R₁ is an alkyl group (which may be cyclic, branched, or linear), typically with from 1 to about 11 carbon atoms, or an arylalkyl group, typically with from 7 to about 18 carbon atoms, R₂ is an alkyl group (and may be either saturated or unsaturated, cyclic, branched, or linear), typically with from 1 to about 11 carbon atoms, or an arylalkyl group, typically with from 7 to about 18 carbon atoms, and R₃ is an alkyl group, including both saturated, unsaturated, linear, branched, and cyclic alkyl groups, preferably with from 1 to about 11 carbon atoms, more preferably with from 1 to about 5 carbon atoms, even more preferably with from 1 to about 3 carbon atoms, and most preferably with 1 carbon atom, or a substituted alkyl group, preferably with from 1 to about 11 carbon atoms, more preferably with from 1 to about 5 carbon atoms, even more preferably with from 1 to about 3 carbon atoms, and most preferably with 1 carbon atom, and X is a halogen atom, such as fluorine, chlorine, bromine, or iodine. Examples of suitable substituents on the substituted alkyl group include (but are not limited to) alkyl groups, including saturated, unsaturated, linear, branched, and cyclic alkyl groups, preferably with from 1 to about 6 carbon atoms, substituted alkyl groups, preferably with from 1 to about 6 carbon atoms, anyl groups, preferably with from 6 to about 24 carbon atoms, substituted anyl groups, preferably with from 6 to about 24 carbon atoms, arylalkyl groups, preferably with from 7 to about 30 carbon atoms, substituted arylalkyl groups, preferably with from 7 to about 30 carbon atoms, alkoxy groups, preferably with from 1 to about 6 carbon atoms, substituted alkoxy groups, preferably with from 1 to about 6 carbon atoms, aryloxy groups, preferably with from 6 to about 24 carbon atoms, substituted aryloxy groups, preferably with from 6 to about 24 carbon atoms, arylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, substituted anylalkyloxy groups, preferably with from 7 to about 30 carbon atoms, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, ester groups, amide groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, mercapto groups, nitroso groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, and the like, wherein the substituents on the substituted alkyl groups, substituted aryl groups, substituted arylalkyl groups, substituted alkoxy groups, substituted aryloxy groups, and substituted arylalkyloxy groups can be (but are not limited to) hydroxy groups, amine groups, imine groups, ammonium groups, pyridine groups, pyridinium groups, ether groups, aldehyde groups, ketone groups, ester groups, amide groups, carboxylic acid groups, carbonyl groups, thiocarbonyl groups, sulfate groups, sulfonate groups, sulfide groups, sulfoxide groups, phosphine groups, phosphonium groups, phosphate groups, cyano groups, nitrile groups, mercapto groups, nitroso groups, halogen atoms, nitro groups, sulfone groups, acyl groups, acid anhydride groups, azide groups, mixtures thereof, and the like, wherein two or more substituents can be joined together to form a ring.

While not being limited to any particular theory, it is believed that thermal cure can also lead to extraction of the

hydroxy group and to crosslinking or chain extension at the "long" bond sites as shown below:

50

45

If desired, the hydroxyalkylated polymer can be further reacted with an unsaturated acid chloride to substitute some or all of the hydroxyalkyl groups with photosensitive groups such as acryloyl or methacryloyl groups or other unsaturated ester groups. The reaction can take place in the presence of triethylamine, which acts as a hydrochloric acid scavenger to form NEt₃H+Cr. Examples of suitable reactants include acryloyl chloride, methacryloyl chloride, cinnamoyl chloride, crotonoyl chloride, ethacryloyl chloride, oleyl chloride, linoleyl chloride, maleoyl chloride, fumaroyl chloride, itaconoyl chloride, citraconoyl chloride, acid chlorides of phenylmaleic acid, 3-hexene-1,6-dicarboxylic acid, and the like. Examples of suitable solvents include 1,1,2,2-tetrachloroethane, methylene chloride, and the like. Typically, the reactants are present in relative amounts with respect to each other by weight of about 1 part hydroxyalkylated polymer, about 1 part triethylamine, about 30 parts solvent, and about 1 part acid chloride.

Some or all of the hydroxyalkyl groups can be replaced with unsaturated ester substituents. Longer reaction times generally lead to greater degrees of substitution of hydroxyalkyl groups with unsaturated ester substituents.

Typical reaction temperatures are from about 0 to about 50°C, and preferably from about 10 to about 25°C, although the temperature can be outside this range. Typical reaction times are from about 1 to about 24 hours, and preferably from about 5 to about 16 hours, although the time can be outside these ranges.

The general reaction scheme is as follows:

45

30

35

40

50

5
$$(CH_{2}OH)_{0} (CH_{2}OH)_{b} (CH_{2}OH)_{c} (CH_{2}OH)_{d}$$

$$H_{2}C = CH - C' C_{1} C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{3}H_{5}$$

$$C_{4}H_{5}$$

$$C_{5}H_{5}$$

$$C_{7}H_{7}$$

$$C_{7}H_$$

wherein a, b, c, d, e, f, g, h, i, j, k, and m are each integers of 0, 1, 2, 3, or 4, provided that the sum of i+e is no greater than 4, the sum of j+f is no greater than 4, the sum of k+g is no greater than 4, and the sum of m+h is no greater than 4, provided that at least one of a, b, c, and d is equal to or greater than 1 in at least some of the monomer repeat units of the polymer, and provided that at least one of e, f, g, and h is equal to at least 1 in at least some of the monomer repeat units of the polymer, and n is an integer representing the number of repeating monomer units. In the corresponding reaction with methacryloyl chloride, the

$$C$$
-CH=CH₂

groups are replaced with

55

45

groups.

5

10

15

20

25

Higher degrees of hydroxyalkylation generally lead to higher degrees of substitution with unsaturated ester groups and thereby to greater photosensitivity of the polymer. Different degrees of substitution may be desirable for different applications. Too high a degree of substitution may lead to excessive sensitivity, resulting in crosslinking or chain extension of both exposed and unexposed polymer material when the material is exposed imagewise to activating radiation, whereas too low a degree of substitution may be undesirable because of resulting unnecessarily long exposure times or unnecessarily high exposure energies. For applications wherein the photopatternable polymer is to be used as a layer in a thermal ink jet printhead, the degree of acryloylation (i.e., the average number of unsaturated ester groups per monomer repeat unit) preferably is from about 0.5 to about 1.2, and more preferably from about 0.65 to about 0.8, although the degree of substitution can be outside these ranges for ink jet printhead applications. Optimum amounts of unsaturated ester substitution are from about 0.8 to about 1.3 milliequivalents of unsaturated ester group per gram of resin.

Some or all of the hydroxyalkyl groups can be replaced with unsaturated ester substituents. Longer reaction times generally lead to greater degrees of substitution of hydroxyalkyl groups with unsaturated ester substituents.

While not being limited to any particular theory, it is believed that exposure to, for example, ultraviolet radiation generally leads to crosslinking or chain extension at the "long" bond sites as shown below for the specific example of polymers having acryloyl functional groups, wherein the ethylenic linkage in the acryloyl group is opened to form the link:

Many of the photosensitivity-imparting groups which are indicated above as being capable of enabling crosslinking or chain extension of the polymer upon exposure to actinic radiation can also enable crosslinking or chain extension of the polymer upon exposure to elevated temperatures; thus the polymers of the present invention can also, if desired, be used in applications wherein thermal curing is employed.

In all of the above reactions and substitutions illustrated above for the polymer of the formula

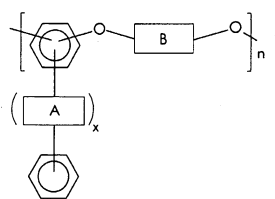
it is to be understood that analogous reactions and substitutions will occur for the polymer of the formula

10

15

20

5



25

30

35

45

50

Photopatternable polymers prepared by the process of the present invention can be used as components in ink jet printheads. The printheads of the present invention can be of any suitable configuration. An example of a suitable configuration, suitable in this instance for thermal ink jet printing, is illustrated schematically in Figure 1, which depicts an enlarged, schematic isometric view of the front face 29 of a printhead 10 showing the array of droplet emitting nozzles 27. Referring also to Figure 2, discussed later, the lower electrically insulating substrate or heating element plate 28 has the heating elements 34 and addressing electrodes 33 patterned on surface 30 thereof, while the upper substrate or channel plate 31 has parallel grooves 20 which extend in one direction and penetrate through the upper substrate front face edge 29. The other end of grooves 20 terminate at slanted wall 21, the floor 41 of the internal recess 24 which is used as the ink supply manifold for the capillary filled ink channels 20, has an opening 25 therethrough for use as an ink fill hole. The surface of the channel plate with the grooves are aligned and bonded to the heater plate 28, so that a respective one of the plurality of heating elements 34 is positioned in each channel, formed by the grooves and the lower substrate or heater plate. Ink enters the manifold formed by the recess 24 and the lower substrate 28 through the fill hole 25 and by capillary action, fills the channels 20 by flowing through an elongated recess 38 formed in the thick film insulative layer 18. The ink at each nozzle forms a meniscus, the surface tension of which prevents the ink from weeping therefrom. The addressing electrodes 33 on the lower substrate or channel plate 28 terminate at terminals 32. The upper substrate or channel plate 31 is smaller than that of the lower substrate in order that the electrode terminals 32 are exposed and available for wire bonding to the electrodes on the daughter board 19, on which the printhead 10 is permanently mounted. Layer 18 is a thick film passivation layer, discussed later, sandwiched between the upper and lower substrates. This layer is etched to expose the heating elements, thus placing them in a pit, and is etched to form the elongated recess to enable ink flow between the manifold 24 and the ink channels 20. In addition, the thick film insulative layer is etched to expose the electrode terminals.

A cross sectional view of Figure 1 is taken along view line 2-2 through one channel and shown as Figure 2 to show how the ink flows from the manifold 24 and around the end 21 of the groove 20 as depicted by arrow 23. As is disclosed in U.S. Patent 4,638,337, U.S. Patent 4,601,777, and U.S. Patent Re. 32,572, a plurality of sets of bubble generating heating elements 34 and their addressing electrodes 33 can be patterned on the polished surface of a single side polished (100) silicon wafer. Prior to patterning, the multiple sets of printhead electrodes 33, the resistive material that serves as the heating elements 34, and the common return 35, the polished surface of the wafer is coated with an underglaze layer 39 such as silicon dioxide, having a typical thickness of from about 500nm (5,000 Angstroms) to about 2 micrometers (microns), although the thickness can be outside this range. The resistive material can be a doped polycrystalline silicon, which can be deposited by chemical vapor deposition (CVD) or any other well known resistive material such as zirconium boride (ZrB2). The common return and the addressing electrodes are typically aluminum leads deposited on the underglaze and over the edges of the heating elements. The common return ends or terminals 37 and addressing electrode terminals 32 are positioned at predetermined locations to allow clearance for wire bonding

to the electrodes (not shown) of the daughter board 19, after the channel plate 31 is attached to make a printhead. The common return 35 and the addressing electrodes 33 are deposited to a thickness typically of from about 0.5 to about 3 micrometers (microns), although the thickness can be outside this range, with the preferred thickness being 1.5 micrometers (microns).

If polysilicon heating elements are used, they may be subsequently oxidized in steam or oxygen at a relatively high temperature, typically about 1,100°C although the temperature can be above or below this value, for a period of time typically of from about 50 to about 80 minutes, although the time period can be outside this range, prior to the deposition of the aluminum leads, in order to convert a small portion of the polysilicon to SiO₂. In such cases, the heating elements are thermally oxidized to achieve an overglaze (not shown) of SiO₂ with a thickness typically of from about 500 Angstroms to about 1 micron, although the thickness can be outside this range, which has good integrity with substantially no pinholes.

In one embodiment, polysilicon heating elements are used and an optional silicon dioxide thermal oxide layer 17 is grown from the polysilicon in high temperature steam. The thermal oxide layer is typically grown to a thickness of from about 0.5 to about 1 micrometer (micron), although the thickness can be outside this range, to protect and insulate the heating elements from the conductive ink. The thermal oxide is removed at the edges of the polysilicon heating elements for attachment of the addressing electrodes and common return, which are then patterned and deposited. If a resistive material such as zirconium boride is used for the heating elements, then other suitable well known insulative materials can be used for the protective layer thereover. Before electrode passivation, a tantalum (Ta) layer (not shown) can be optionally deposited, typically to a thickness of about 1 micron, although the thickness can be above or below this value, on the heating element protective layer 17 for added protection thereof against the cavitational forces generated by the collapsing ink vapor bubbles during printhead operation. The tantalum layer is etched off all but the protective layer 17 directly over the heating elements using, for example, CF₄/O₂ plasma etching. For polysilicon heating elements, the aluminum common return and addressing electrodes typically are deposited on the underglaze layer and over the opposing edges of the polysilicon heating elements which have been cleared of oxide for the attachment of the common return and electrodes.

25

30

35

40

45

50

55

For electrode passivation, a film 16 is deposited over the entire wafer surface, including the plurality of sets of heating elements and addressing electrodes. The passivation film 16 provides an ion barrier which will protect the exposed electrodes from the ink. Examples of suitable ion barrier materials for passivation film 16 include polyimide, plasma nitride, phosphorous doped silicon dioxide, materials disclosed herein as being suitable for insulative layer 18, and the like, as well as any combinations thereof. An effective ion barrier layer is generally achieved when its thickness is from about 100nm (1000 Angstroms) to about 10 microns, although the thickness can be outside this range. In 300 dpi printheads, passivation layer 16 preferably has a thickness of about 3 microns, although the thickness can be above or below this value. In 600 dpi printheads, the thickness of passivation layer 16 preferably is such that the combined thickness of layer 16 and layer 18 is about 25 microns, although the thickness can be above or below this value. The passivation film or layer 16 is etched off of the terminal ends of the common return and addressing electrodes for wire bonding later with the daughter board electrodes. This etching of the silicon dioxide film can be by either the wet or dry etching method. Alternatively, the electrode passivation can be by plasma deposited silicon nitride (Si₃N₄).

Next, a thick film type insulative layer 18, of a polymeric material discussed in further detail herein, is formed on the passivation layer 16, typically having a thickness of from about 10 to about 100 micrometers (microns) and preferably in the range of from about 25 to about 50 micrometers (microns), although the thickness can be outside these ranges. Even more preferably, in 300 dpi printheads, layer 18 preferably has a thickness of about 30 micrometers (microns), and in 600 dpi printheads, layer 18 preferably has a thickness of from about 20 to about 22 micrometers (microns), although other thicknesses can be employed. The insulative layer 18 is photolithographically processed to enable etching and removal of those portions of the layer 18 over each heating element (forming recesses 26), the elongated recess 38 for providing ink passage from the manifold 24 to the ink channels 20, and over each electrode terminal 32, 37. The elongated recess 38 is formed by the removal of this portion of the thick film layer 18. Thus, the passivation layer 16 alone protects the electrodes 33 from exposure to the ink in this elongated recess 38. Optionally, if desired, insulative layer 18 can be applied as a series of thin layers of either similar or different composition. Typically, a thin layer is deposited, photoexposed, partially cured, followed by deposition of the next thin layer, photoexposure, partial curing, and the like. The thin layers constituting thick film insulative layer 18 contain a polymer of the formula indicated hereinabove. In one embodiment of the present invention, a first thin layer is applied to contact layer 16, said first thin layer containing a mixture of a polymer of the formula indicated hereinabove and an epoxy polymer, followed by photoexposure, partial curing, and subsequent application of one or more successive thin layers containing a polymer of the formula indicated hereinabove.

Figure 3 is a similar view to that of Figure 2 with a shallow anisotropically etched groove 40 in the heater plate, which is silicon, prior to formation of the underglaze 39 and patterning of the heating elements 34, electrodes 33 and common return 35. This recess 40 permits the use of only the thick film insulative layer 18 and eliminates the need for the usual electrode passivating layer 16. Since the thick film layer 18 is impervious to water and relatively thick (typically

from about 20 to about 40 micrometers (microns), although the thickness can be outside this range), contamination introduced into the circuitry will be much less than with only the relatively thin passivation layer 16 well known in the art. The heater plate is a fairly hostile environment for integrated circuits. Commercial ink generally entails a low attention to purity. As a result, the active part of the heater plate will be at elevated temperature adjacent to a contaminated aqueous ink solution which undoubtedly abounds with mobile ions. In addition, it is generally desirable to run the heater plate at a voltage of from about 30 to about 50 volts, so that there will be a substantial field present. Thus, the thick film insulative layer 18 provides improved protection for the active devices and provides improved protection, resulting in longer operating lifetime for the heater plate.

When a plurality of lower substrates 28 are produced from a single silicon wafer, at a convenient point after the underglaze is deposited, at least two alignment markings (not shown) preferably are photolithographically produced at predetermined locations on the lower substrates 28 which make up the silicon wafer. These alignment markings are used for alignment of the plurality of upper substrates 31 containing the ink channels. The surface of the single sided wafer containing the plurality of sets of heating elements is bonded to the surface of the wafer containing the plurality of ink channel containing upper substrates subsequent to alignment.

15

25

45

50

As disclosed in U.S. Patent 4,601,777 and U.S. Patent 4,638,337, the channel plate is formed from a two side polished, (100) silicon wafer to produce a plurality of upper substrates 31 for the printhead. After the wafer is chemically cleaned, a pyrolytic CVD silicon nitride layer (not shown) is deposited on both sides. Using conventional photolithography, a via for fill hole 25 for each of the plurality of channel plates 31 and at least two vias for alignment openings (not shown) at predetermined locations are printed on one wafer side. The silicon nitride is plasma etched off of the patterned vias representing the fill holes and alignment openings. A potassium hydroxide (KOH) anisotropic etch can be used to etch the fill holes and alignment openings. In this case, the [111] planes of the (100) wafer typically make an angle of about 54.7 degrees with the surface of the wafer. The fill holes are small square surface patterns, generally of about 20 mils (500 microns) per side, although the dimensions can be above or below this value, and the alignment openings are from about 60 to about 80 mils (1.5 to 3 millimeters) square, although the dimensions can be outside this range. Thus, the alignment openings are etched entirely through the 20 mil (0.5 millimeter) thick wafer, while the fill holes are etched to a terminating apex at about halfway through to three-quarters through the wafer. The relatively small square fill hole is invariant to further size increase with continued etching so that the etching of the alignment openings and fill holes are not significantly time constrained.

Next, the opposite side of the wafer is photolithographically patterned, using the previously etched alignment holes as a reference to form the relatively large rectangular recesses 24 and sets of elongated, parallel channel recesses that will eventually become the ink manifolds and channels of the printheads. The surface 22 of the wafer containing the manifold and channel recesses are portions of the original wafer surface (covered by a silicon nitride layer) on which an adhesive, such as a thermosetting epoxy, will be applied later for bonding it to the substrate containing the plurality of sets of heating elements. The adhesive is applied in a manner such that it does not run or spread into the grooves or other recesses. The alignment markings can be used with, for example, a vacuum chuck mask aligner to align the channel wafer on the heating element and addressing electrode wafer. The two wafers are accurately mated and can be tacked together by partial curing of the adhesive. Alternatively, the heating element and channel wafers can be given precisely diced edges and then manually or automatically aligned in a precision jig. Alignment can also be performed with an infrared aligner-bonder, with an infrared microscope using infrared opaque markings on each wafer to be aligned, or the like. The two wafers can then be cured in an oven or laminator to bond them together permanently. The channel wafer can then be milled to produce individual upper substrates. A final dicing cut, which produces end face 29, opens one end of the elongated groove 20 producing nozzles 27. The other ends of the channel groove 20 remain closed by end 21. However, the alignment and bonding of the channel plate to the heater plate places the ends 21 of channels 20 directly over elongated recess 38 in the thick film insulative layer 18 as shown in Figure 2 or directly above the recess 40 as shown in Figure 3 enabling the flow of ink into the channels from the manifold as depicted by arrows 23. The plurality of individual printheads produced by the final dicing are bonded to the daughter board and the printhead electrode terminals are wire bonded to the daughter board electrodes.

In one embodiment, a heater wafer with a phosphosilicate glass layer is spin coated with a solution of Z6020 adhesion promoter (0.01 weight percent in 95 parts methanol and 5 parts water, Dow Corning) at 3000 revolutions per minute for 10 seconds and dried at 100°C for between 2 and 10 minutes. The wafer is then allowed to cool at 25°C for 5 minutes before spin coating the photoresist containing the photopatternable polymer onto the wafer at between 1,000 and 3,000 revolutions per minute for between 30 and 60 seconds. The photoresist solution is made by dissolving polyarylene ether ketone with 0.75 acryloyl groups and 0.75 hydroxymethyl groups per repeat unit and a weight average molecular weight of 25,000 in N-methylpyrrolidinone at 40 weight percent solids with Michler's ketone (1.2 parts ketone per every 10 parts of 40 weight percent solids polymer solution). The film is heated (soft baked) in an oven for between 10 and 15 minutes at 70°C. After cooling to 25°C over 5 minutes, the film is covered with a mask and exposed to 365 nanometer ultraviolet light, amounting to between 150 and 1,500 milliJoules per cm². The exposed wafer is then heated at 70°C for 2 minutes post exposure bake, followed by cooling to 25°C over 5 minutes. The film is developed with 60:

40 chloroform/cyclohexanone developer, washed with 90:10 hexanes/cyclohexanone, and then dried at 70°C for 2 minutes. A second developer/wash cycle is carried out if necessary to obtain a wafer with clean features. The processed wafer is transferred to an oven at 25°C, and the oven temperature is raised from 25 to 90°C at 2°C per minute. The temperature is maintained at 90°C for 2 hours, and then increased to 260°C at 2°C per minute. The oven temperature is maintained at 260°C for 2 hours and then the oven is turned off and the temperature is allowed to cool gradually to 25°C. When thermal cure of the photoresist films is carried out under inert atmosphere, such as nitrogen or one of the noble gases, such as argon, neon, krypton, xenon, or the like, there is markedly reduced oxidation of the developed film and improved thermal and hydrolytic stability of the resultant devices. Moreover, adhesion of developed photoresist film is improved to the underlying substrate. If a second layer is spin coated over the first layer, the heat cure of the first developed layer can be stopped between 80 and 260°C before the second layer is spin coated onto the first layer. A second thicker layer is deposited by repeating the above procedure a second time. This process is intended to be a guide in that procedures can be outside the specified conditions depending on film thickness and photoresist molecular weight. Films at 30 micrometers (microns) have been developed with clean features at 600 dots per inch.

The present invention also encompasses printing processes with printheads according to the present invention. One embodiment of the present invention is directed to an ink jet printing process which comprises (1) preparing an ink jet printhead comprising a plurality of channels, wherein the channels are capable of being filled with ink from an ink supply and wherein the channels terminate in nozzles on one surface of the printhead, said preparation being according to the process of the present invention; (2) filling the channels with an ink; and (3) causing droplets of ink to be expelled from the nozzles onto a receiver sheet in an image pattern. A specific embodiment of this process is directed to a thermal ink jet printing process, wherein the droplets of ink are caused to be expelled from the nozzles by heating selected channels in an image pattern. The droplets can be expelled onto any suitable receiver sheet, such as fabric, plain paper such as Xerox® 4024 or 4010, coated papers, or transparency materials.

All parts and percentages in the Examples are by weight unless otherwise indicated.

EXAMPLE I

5

10

25

40

45

50

55

A polyarylene ether ketone of the formula

30 H₃C CH₃

wherein n is between about 2 and about 30 (hereinafter referred to as poly(4-CPK-BPA)) was prepared as follows. A 5-liter, 3-neck round-bottom flask equipped with a Dean-Stark (Barrett) trap, condenser, mechanical stirrer, argon inlet, and stopper was situated in a silicone oil bath. 4,4'-Dichlorobenzophenone (Aldrich 11,370, Aldrich Chemical Co., Milwaukee, WI, 250 grams), bis-phenol A (Aldrich 23,965-8, 244.8 grams), potassium carbonate (327.8 grams), anhydrous *N,N*-dimethylacetamide (1,500 milliliters), and toluene (275 milliliters) were added to the flask and heated to 175°C (oil bath temperature) while the volatile toluene component was collected and removed. After hours of heating 30 hours at 175°C with continuous stirring, the reaction mixture was filtered to remove insoluble salts, and the resultant solution was added to methanol (5 gallons) to precipitate the polymer. The polymer was isolated by filtration, and the wet filter cake was washed with water (3 gallons) and then with methanol (3 gallons). The yield was 360 grams of vacuum dried polymer. The molecular weight of the polymer was determined by gel permeation chromatography (gpc) (elution solvent was tetrahydrofuran) with the following results: M_n 2,800, M_{peak} 5,800, M_w 6,500, M_z 12,000 and M_{z+1} 17,700. As a result of the stoichiometries used in the reaction, it is believed that this polymer had end groups derived from bis-phenol A. When the reaction was allowed to proceed for 35, 40, and 48 hours at 175°C, the respective values of M_n of the poly(4-CPK-BPA) formed were 3,000, 3,300, and 4,000.

A solution containing 100 parts by weight of the polyarylene ether ketone thus prepared having a M_n of 2,800, 44.5 parts by weight of paraformaldehyde, 1 part by weight sodium hydroxide, and 1 part by weight tetramethylammonium hydroxide in 200 parts by weight 1,1,2,2-tetrachloroethane was heated at 100°C. Vigorous stirring and heating were continued for 16 hours. The resultant mixture was extracted with water and the organic layer was dried over magnesium sulfate. After precipitation into methanol, the filtered polymer was vacuum dried to yield 100 parts by weight hydroxymethylated polyarylene ether ketone with 1.0 hydroxymethyl group per repeat unit.

Thereafter, 1 part by weight of the hydroxymethylated polymer thus formed was allowed to react with 1 part by weight of isocyanatoethyl methacrylate in 20 parts by weight methylene chloride to form an acryloylated and hydroxymethylated polymer at 25°C within 16 hours. The resultant polymer had about 0.7 acryloyl groups per repeat unit.

EXAMPLE II

5

10

15

20

A hydroxymethylated polyarylene ether ketone was prepared as described in Example I. One part by weight of the hydroxymethylated polymer was allowed to react with 1 part by weight of acryloyl chloride in 30 parts by weight methylene chloride in the presence of 1 part by weight triethylamine. The reaction mixture was cooled in an ice bath, and the ice bath was allowed to melt while the reaction mixture was stirred at 25°C for 16 hours. The resultant polymer had 0.6 acryloyl groups per repeat unit.

EXAMPLE III

A polyarylene ether ketone of the formula

25

30

35

40

wherein n is between about 2 and about 30 (hereinafter referred to as poly(4-CPK-BPA)) was prepared as follows. A 5 liter, 3-neck round-bottom flask equipped with a Dean-Stark (Barrett) trap, condenser, mechanical stirrer, argon inlet, and stopper was situated in a silicone oil bath. 4,4'-Dichlorobenzophenone (Aldrich 11,370, Aldrich Chemical Co., Milwaukee, WI, 250 grams), bis-phenol A (Aldrich 23,965-8, 244.8 grams), potassium carbonate (327.8 grams), anhydrous N,N-dimethylacetamide (1,500 milliliters), and toluene (275 milliliters) were added to the flask and heated to 175°C (oil bath temperature) while the volatile toluene component was collected and removed. After 48 hours of heating at 175°C with continuous stirring, the reaction mixture was filtered to remove insoluble salts, and the resultant solution was added to methanol (5 gallons) to precipitate the polymer. The polymer was isolated by filtration, and the wet filter cake was washed with water (3 gallons) and then with methanol (3 gallons). The yield was 360 grams of vacuum dried polymer. The molecular weight of the polymer was determined by gel permeation chromatography (gpc) (elution solvent was tetrahydrofuran) with the following results: M_n 3,430, M_{peak} 5,380, M_w 3,600, M_z 8,700, and M_{z+1} 12,950. The glass transition temperature of the polymer was between 125 and 155°C as determined using differential scanning calorimetry at a heating rate of 20°C per minute dependent on molecular weight. Solution cast films from methylene chloride were clear, tough, and flexible. As a result of the stoichiometries used in the reaction, it is believed that this polymer had end groups derived from bis-phenol A.

45

A solution of chloromethyl ether in methyl acetate was made by adding 282.68 grams (256 milliliters) of acetyl chloride to a mixture of dimethoxy methane (313.6 grams, 366.8 milliliters) and methanol (10 milliliters) in a 5 liter 3-neck round-bottom flask equipped with a mechanical stirrer, argon inlet, reflux condenser, and addition funnel. The solution was diluted with 1,066.8 milliliters of 1,1,2,2-tetrachloroethane and then tin tetrachloride (2.4 milliliters) was added via a gas-tight syringe, along with 1,1,2,2-tetrachloroethane (133.2 milliliters) using an addition funnel. The reaction solution was heated to 50°C and a solution of poly(4-CPK-BPA) (160.8 grams) in 1,1,2,2-tetrachloroethane (1,000 milliliters) was rapidly added. The reaction mixture was then heated to reflux with an oil bath set at 110°C. After four hours reflux with continuous stirring, heating was discontinued and the mixture was allowed to cool to 25°C. The reaction mixture was transferred in stages to a 2 liter round bottom flask and concentrated using a rotary evaporator with gentle heating up to 50°C and reduced pressure maintained with a vacuum pump trapped with liquid nitrogen. The concentrate was added to methanol (6 gallons) to precipitate the polymer using a Waring blender. The polymer was isolated by filtration and vacuum dried to yield 200 grams of poly(4-CPK-BPA) with 1.5 chloromethyl groups per repeat unit as identified using ¹H NMR spectrometry.

55

50

Claims

A composition which comprises (a) a polymer containing at least some monomer repeat units with photosensitivity-

imparting substituents which enable crosslinking or chain extension of the polymer upon exposure to actinic radiation, said polymer being of the formula

A X O B

10 or

5

35

55

20 B O B

wherein x is an integer of 0 or 1, A is

30 <u>C</u> C

40

45 OSSO

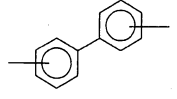
50 OI

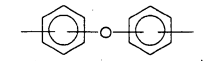
-C(CH₃)₂-

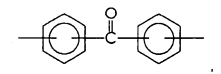
-0-

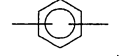
or mixtures thereof, B is

5 H₃C, C₂H₅

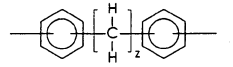




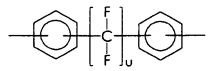




wherein v is an integer of from 1 to about 20,



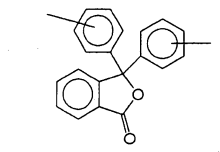
wherein z is an integer of from 2 to about 20,

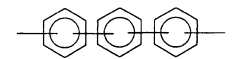


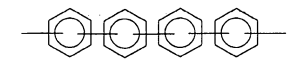
wherein u is an integer of from 1 to about 20,

wherein w is an integer of from 1 to about 20,

ÇH₃ H₃C

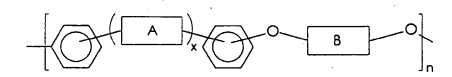




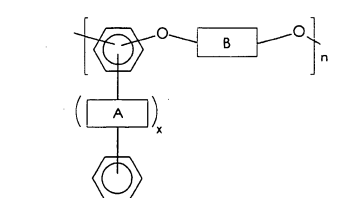


or mixtures thereof, and n is an integer representing the number of repeating monomer units, wherein said photosensitivity-imparting substituents are hydroxyalkyl groups; (b) at least one member selected from the group consisting of photoinitiators and sensitizers; and (c) an optional solvent.

2. A composition which comprises a crosslinked or chain extended polymer of the formula



or



wherein x is an integer of 0 or 1, A is

.

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ &$$

-C(CH₃)₂-,

-O-,

or mixtures thereof, B is

5

10

15

20

25

45

50

30 H₃C, CH₃

H₃C₂H₅

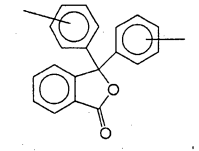
wherein v is an integer of from 1 to about 20,

wherein z is an integer of from 2 to about 20,

wherein u is an integer of from 1 to about 20,

wherein w is an integer of from 1 to about 20,

CH₃
CH₃
CH₃



$$-000$$



or mixtures thereof, and n is an integer representing the number of repeating monomer units, at least some of said crosslinking or chain extension occuring through groups of the formula

5 NH NH

wherein R is

-c-

-R1C=CR2-,

25 or

15

20

30

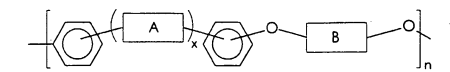
40

45

R₃ —C— R₄

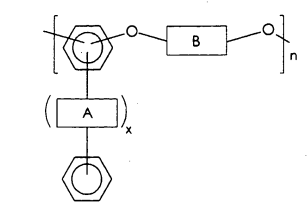
wherein R₁, R₂, R₃, and R₄ each, independently of the others, is a hydrogen atom, an alkyl group, a substituted aryl group, an arylalkyl group, or a substituted arylalkyl group, and x is 0 or 1.

3. A composition which comprises a crosslinked or chain extended polymer of the formula



or

50



wherein x is an integer of 0 or 1, A is

30 C

-O-,

-C(CH₃)₂-,

or mixtures thereof, B is

5

10

15

20

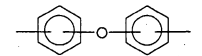
25

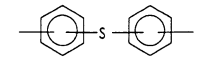
45

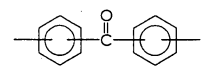
50

F₃C, CF₃

H₃C_{...}C₂H₅





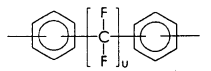






wherein v is an integer of from 1 to about 20,

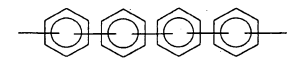
wherein z is an integer of from 2 to about 20,



wherein u is an integer of from 1 to about 20,

CH₃ CH₃

wherein w is an integer of from 1 to about 20,



or mixtures thereof, and n is an integer representing the number of repeating monomer units, at least some of said crosslinking or chain extension occuring through groups of the formula

wherein R_1 is an alkyl group or an arylalkyl group, R_2 is an alkyl group or an arylalkyl group, and R_3 is an alkyl group or a substituted alkyl group.

- 4. A composition according to any of claims 1 to 3 wherein the polymer has end groups derived from the "A" groups of the polymer.
 - A composition according to any of claims 1 to 3 wherein the polymer has end groups derived from the "B" groups of the polymer.
- 35 6. A process for forming a thermal ink jet printhead comprising the steps of:

5

10

15

20

25

40

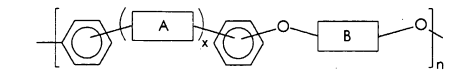
45

50

- (a) depositing a layer (18) comprising the polymer-containing composition onto a lower substrate (28) in which one surface thereof has an array of heating elements (34) and addressing electrodes (33) having terminal ends (32) formed thereon;
- (b) exposing the layer (18) to actinic radiation in an imagewise pattern such that the polymer in exposed areas becomes crosslinked or chain extended and the polymer in unexposed areas does not become crosslinked or chain extended, wherein the unexposed areas correspond to areas of the lower substrate (28) having thereon the heating elements (34) and the terminal ends (32) of the addressing electrodes (33);
- (c) removing the polymer from the unexposed areas, thereby forming recesses in the layer (18), said recesses exposing the heating elements (34) and the terminal ends (32) of the addressing electrodes (33);
- (d) providing an upper substrate (31) with a set of parallel grooves (20) for subsequent use as ink channels and a recess for subsequent use as a manifold, the grooves (20) being open at one end for serving as droplet emitting nozzles; and
- (e) aligning, mating, and bonding the upper (31) and lower (28) substrates together to form a printhead with the grooves (20) in the upper substrate (31) being aligned with the heating elements (34) in the lower substrate (28) to form droplet emitting nozzles, thereby forming a thermal ink jet printhead.
- 7. An ink jet printhead (10) which comprises (i) an upper substrate (31) with a set of parallel grooves (20) for subsequent use as ink channels and a recess (24) for subsequent use as a manifold, the grooves (20) being open at one end for serving as droplet emitting nozzles, (ii) a lower substrate (28) in which one surface thereof has an array of heating elements (34) and addressing electrodes (33) formed thereon, and (iii) a layer (18) deposited on the surface of the lower substrate (28) and over the heating elements (34) and addressing electrodes (33) and patterned to form recesses therethrough to expose the heating elements (34) and terminal ends (32) of the ad-

dressing electrodes (33), the upper (31) and lower substrates (28) being aligned, mated, and bonded together to form the printhead (10) with the grooves (20) in the upper substrate (31) being aligned with the heating elements (34) in the lower substrate (28) to form droplet emitting nozzles, said layer (18) comprising a polymer-containing composition according to any of claims 1 to 5.

8. A process which comprises the steps of: (a) reacting a haloalkylated polymer of the formula



or

5

10

15

25 (A)_X

wherein x is an integer of 0 or 1, A is

35 C

45

50 SS

-O-,

-C(CH₃)₂-,

or mixtures thereof, B is

F₃C₂ CF₃



wherein v is an integer of from 1 to about 20,

(CH₂

wherein z is an integer of from 2 to about 20,

wherein u is an integer of from 1 to about 20,

wherein w is an integer of from 1 to about 20,





20



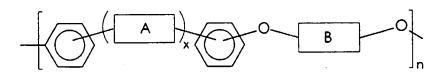


or mixtures thereof, and n is an integer representing the number of repeating monomer units, with water and a base, thereby forming a polymer with hydroxyalkyl groups; and optionally (b) converting the hydroxyalkyl groups 30 to unsaturated ester groups.

9. A process which comprises the steps of: (a) reacting a polymer of the formula

35

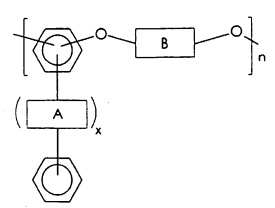
40

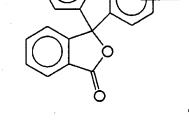


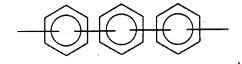
or

45

50







.wherein x is an integer of 0 or 1, A is

•

-C(CH₃)₂-,

-O-,

or mixtures thereof, B is

5

10

15

20

25

35

40

H₃C, CH₃

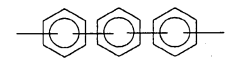
5 wherein v is an integer of from 1 to about 20,

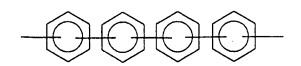
wherein z is an integer of from 2 to about 20,

wherein u is an integer of from 1 to about 20,

wherein w is an integer of from 1 to about 20,

15 CH₃





or mixtures thereof, and n is an integer representing the number of repeating monomer units, with formaldehyde or paraformaldehyde and a base, thereby forming a photopatternable polymer with hydroxymethyl groups; and optionally (b) converting the hydroxymethyl groups to unsaturated ester groups.

10. A process according to either of claims 8 or 9 wherein crosslinking or chain extension is effected by heating the polymer to a temperature sufficient to enable the photosensitivity-imparting groups to form crosslinks or chain extensions in the polymer, or by exposing the polymer to actinic radiation such that the polymer in exposed areas becomes crosslinked or chain extended.

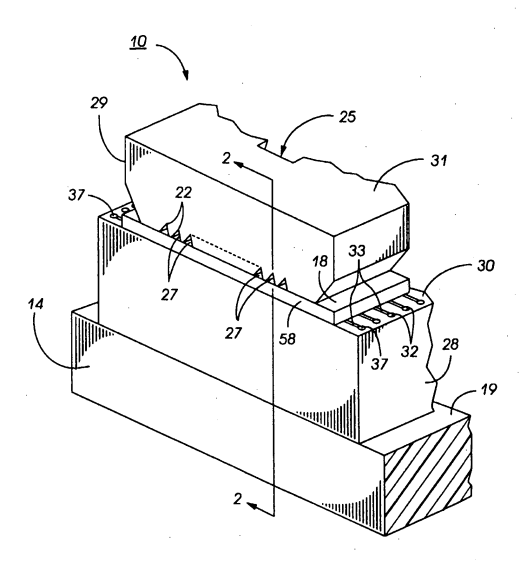


FIG. 1

